

# Design Requirements for Interfering Particles To Maintain Coadaptive Stability with HIV-1

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Defective interfering particles (DIPs) are viral deletion mutants lacking essential transacting or packaging elements and must be complemented by wild-type virus to propagate. DIPs transmit through human populations, replicating at the expense of the wild-type virus and acting as molecular parasites of viruses. Consequently, engineered DIPs have been proposed as therapies for a number of diseases, including human immunodeficiency virus (HIV). However, it is not clear if DIP-based therapies would face evolutionary blocks given the high mutation rates and high within-host diversity of lentiviruses. Divergent evolution of HIV and DIPs appears likely since natural DIPs have not been detected for lentiviruses, despite extensive sequencing of HIVs and simian immunodeficiency viruses (SIVs). Here, we tested if the apparent lack of lentiviral DIPs is due to natural selection and analyzed which molecular characteristics a DIP or DIP-based therapy would need to maintain coadaptive stability with HIV-1. Using a well-established mathematical model of HIV-1 in a host extended to include its replication in a single cell and interference from DIP, we calculated evolutionary selection coefficients. The analysis predicts that interference by codimerization between DIPs and HIV-1 genomes is evolutionarily unstable, indicating that recombination between DIPs and HIV-1 would be selected against. In contrast, DIPs that interfere via competition for capsids have the potential to be evolutionarily stable if the capsid-to-genome production ratio of HIV-1 is  $>1$ . Thus, HIV-1 variants that attempt to “starve” DIPs to escape interference would be selected against. In summary, the analysis suggests specific experimental measurements that could address the apparent lack of naturally occurring lentiviral DIPs and specifies how therapeutic approaches based on engineered DIPs could be evolutionarily robust and avoid recombination.

**D**efective interfering particles (DIPs) are mutant versions of viruses that contain significant genomic deletions such that they are unable to replicate except when complemented by wild-type virus replicating within the same cell (for reviews, see references 1 and 2). At the most fundamental level, DIPs arise because viral genomes encode both *cis*- and *trans*-acting elements. *Trans*-acting elements (*trans* elements) code for gene products, such as capsid proteins or transcription factors, and *cis*-acting elements (*cis* elements) are regions of the viral genome that interact with *trans*-element products to achieve productive viral replication, including viral genome amplification, encapsidation, and viral egress. In the case of RNA virus genomes, *cis* elements can include viral enhancers and promoters and also viral genome-packaging signals. Viral capsid and envelope proteins, on the other hand, are examples of *trans* elements. Mutations that result in the loss of at least one obligate *trans* element but retain all necessary *cis* elements required for productive replication can generate DIPs.

DIPs have been reported for several important human and animal pathogens, including murine leukemia virus (3), Rous sarcoma virus (4), vesicular stomatitis virus (5), influenza virus (6), and dengue fever virus (7,8). Presumably, the characteristic error-prone replication of RNA viruses leads to frequent generation of defective mutants (1,2). In some cases (e.g., murine leukemia virus), specific mutations within DIPs encode novel *trans* elements that enhance immune responses to infected cells or produce cytoidal or cytotoxic products that lead to enhanced virulence (3,9–11). Many DIPs interfere with the *in vivo* replication of the wild-type pathogen and attenuate virulence and reduce disease in animal models (12–16). Based on these findings, therapies

based on DIPs have been proposed for a number of diseases, including human immunodeficiency virus type 1 (HIV-1) (15,17).

Here, we considered a broad class of DIP designs that represent HIV-1 with mutations or deletions in capsid proteins and Nef responsible for protection against superinfection. Possible designs range from minimal DIPs completely lacking any *trans* elements to DIPs expressing some *trans* elements, including transcriptional transactivating proteins (Tat) and genomic export proteins (Rev). All these HIV-1 mutants can be expressed, complemented, and packaged by either providing the missing viral proteins in *trans* (18) or coinfecting with a homologous replication-competent virus (19) (e.g., wild-type HIV or a “helper” virus) which provides the missing *trans* factors. This shunting of *trans* factors to the DIP interferes with the intracellular replication of the wild-type virus. Since DIPs coopt viral replication or packaging machinery and transmit their genetic material at the expense of the wild-type virus, they are essentially molecular parasites of wild-type viruses. However, whether coevolution might occur between HIV-1 and a DIP therapy and whether HIV-1 could escape from a DIP have not been explored.

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Importantly, natural lentiviral DIPs have not been detected, despite extensive sequencing of HIV and simian immunodeficiency virus (SIV) strains. For example, although many cells infected with HIV-1 harbor permanently or transiently inactive variants of HIV-1 provirus, the vast majority of infected cells contain a single provirus (20). There is no evidence in humans or nonhuman primates of coinfection of cells with a replication-competent HIV-1 (or SIV) provirus and a replication-deficient HIV-1 (or SIV) mutant that is mobilized by a replication-competent virus and could be identified as a DIP. This absence of lentiviral DIPs could result from one of three possibilities: (i) a lack of assays sensitive enough to detect existing DIPs, (ii) molecular restrictions resulting from the unique biology of lentiviruses compared to other RNA viruses (i.e., a “molecular block” to DIP formation), or (iii) rapid evolutionary selection against DIPs that do spontaneously arise (i.e., lentiviral DIPs could be generated with appreciable frequency but are evolutionarily unstable and exist only transiently before being “selected out” of the viral population, either within an infected individual or at the level of the infected host population in a “boom-then-bust” scenario) (see Discussion).

A molecular block to DIP formation is unlikely in the case of HIV-1. While different viruses and target cells clearly vary in their ability to form and propagate DIPs (21–23), there do not appear to be any inherent blocks to the generation of large deletions within the HIV-1 genome (24) or in the packaging or mobilization of subgenomic HIV-1. Subgenomic HIV-1 strains are routinely produced from packaging cell lines during the generation of lentiviral vectors for gene delivery and gene therapy (18). Moreover, numerous studies have artificially engineered HIV-1 variants that lack *trans* elements but conditionally replicate in the presence of full-length HIV-1 and transmit through tissue cultures (25–30). A conditionally replicating HIV-1 construct termed VRX496, which is essentially a DIP encoding an antisense gene therapy element, has even been reported to mobilize within a humanized-mouse model of HIV-1 (31) and in patients infected with HIV-1 (15).

While a molecular block to lentiviral DIPs appears unlikely, it is unclear if DIPs are evolutionarily stable, since HIV-1 might escape by mutation. While studies have quantitatively explored the dynamics and dynamic stability (i.e., conditions for survival) of DIPs (17, 32–35), there have been no quantitative analyses of the long-term evolutionary stability of lentiviral DIPs. Experimentally, DIPs for vesicular stomatitis virus are known to exert strong selective pressures on their helper virus as a result of the divestment of cellular resources to the DIPs during intracellular replication (i.e., DIP-mediated resource stealing). In long-term cultures of cells infected with vesicular stomatitis virus and its DIP (T particle) over a period of 5 years and several hundred passages, Holland and coworkers observed significant coevolution of both wild-type virus and a DIP (23, 36), including the emergence of formerly susceptible wild-type virus that gained resistance to the DIP from previous passages (21, 37–39).

One would expect HIV-1 to be under pressure to change genetically to minimize its interaction with a DIP. Since HIV-1 demonstrates a considerable degree of evolutionary plasticity, it is likely to eventually escape the parasite. However, evolutionary plasticity of the DIPs would be similar to that of HIV-1 and under similar selective pressure to maintain the parasitization of HIV-1. This would establish a coevolutionary arms race between HIV-1

and the DIP, and the outcome of such a race would not be obvious, especially when the system is further complicated by dynamics at multiple scales of biological organization (e.g., the single-cell and individual levels), as is the case for HIV-1 and the DIP.

The appropriate tools for theoretically analyzing such complex coevolutionary processes are found in the field of population genetics. Simple models of HIV-1 population genetics, introduced in the 1990s, proved to be powerful tools for the analysis of HIV-1 sequence data and led to predictive experimentally tested theories on the rates of HIV-1 adaptation and selection and its effective recombination rate (40–50). Most relevantly, population genetics models estimated the fitness costs of HIV-1 mutations (including drug resistance and immune escape mutations) by calculating evolutionary selection coefficients from the data (40, 43, 46, 48, 51–54). Coupled with the well-established models of HIV-1 dynamics in individuals (55, 56), population genetics provides a rigorous approach to studying the coevolution of HIV-1 and DIPs and to addressing the question of whether lentiviral DIPs would be evolutionarily stable.

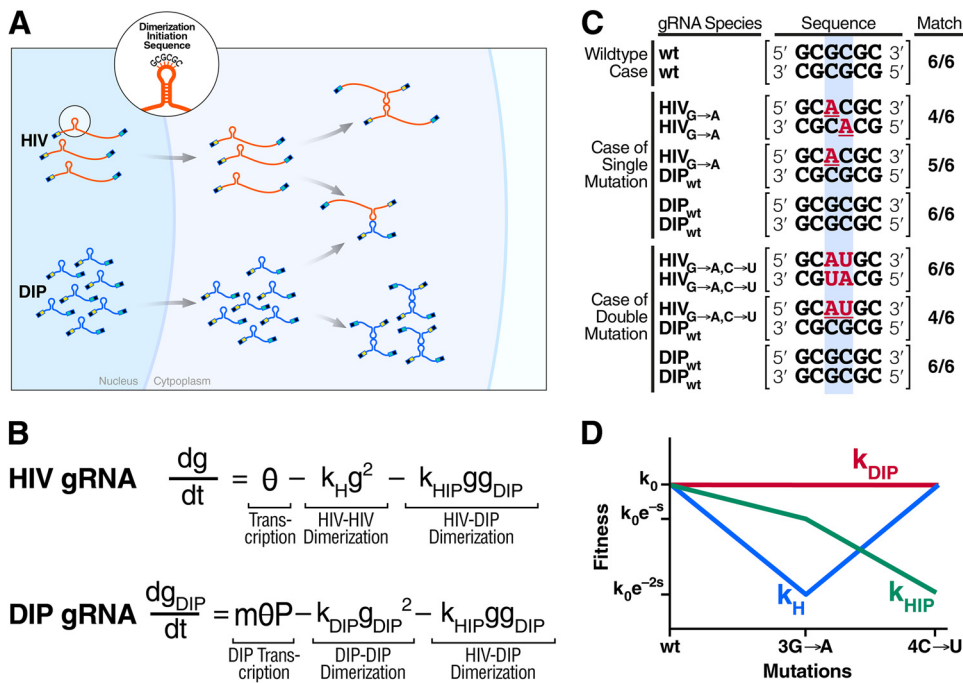
In this study, we calculated the likely direction of HIV-1 and DIP evolution within an infected individual to determine if DIP-based therapies would face an evolutionary disadvantage. Specifically, we asked whether two mechanisms of competition corresponding to two points in the viral replication cycle—*cis* stealing (e.g., competition between single-stranded HIV genomic RNA and single-stranded DIP RNA to bind a second strand of HIV genomic RNA) and *trans* stealing (e.g., competition for viral capsids)—are evolutionarily stable strategies of interference. The practical aim of this inquiry was to direct the design of future DIP vectors. Testing stability in cell culture is not a trivial matter, because the total number of cells, even in large flasks, is several logs lower than in an animal/human host, and the speed of evolution may be sensitive to rare double or triple mutations. Therefore, before attempting expensive long-term animal studies, it was important to test evolutionary stability *in silico*.

To do this, we considered a model of HIV-DIP interaction at two scales: (i) on the single-cell scale, a simplified intracellular model approximated the molecular mechanisms of HIV-1 particle formation and DIP formation (two models for two types of interference), and (ii) on the individual-patient scale, a standard model of HIV-1 dynamics in individuals (55, 56) was extended to include the presence of DIPs. We derived the fitness effect of mutation at the within-host scale (effective selection coefficient) directly from the underlying intracellular model and tested whether HIV-1 is likely to evolve to decrease DIP-mediated stealing of its products and hence escape suppression by the DIP.

## THEORY

**Preliminary analysis: DIP interference by the competition for HIV-1 genomic RNA leads to divergent evolution at the single-cell level.** We begin by analyzing the stability of DIPs that interfere via binding to the genomic RNAs (gRNAs) of the wild-type lentivirus (i.e., genome stealing). Lentiviruses are diploid, and gRNAs are packaged into virions in pairs, where encapsidation of two copies of RNA is achieved by allowing the gRNAs to dimerize. This gRNA pairing is initiated at a six-nucleotide palindrome termed the dimerization initiation signal (DIS), which is located within stem-loop 1 (SL1) of the HIV-1 genome and has the consensus sequence GCGCGC (57).

A minimal mathematical model that considers only nondimerized



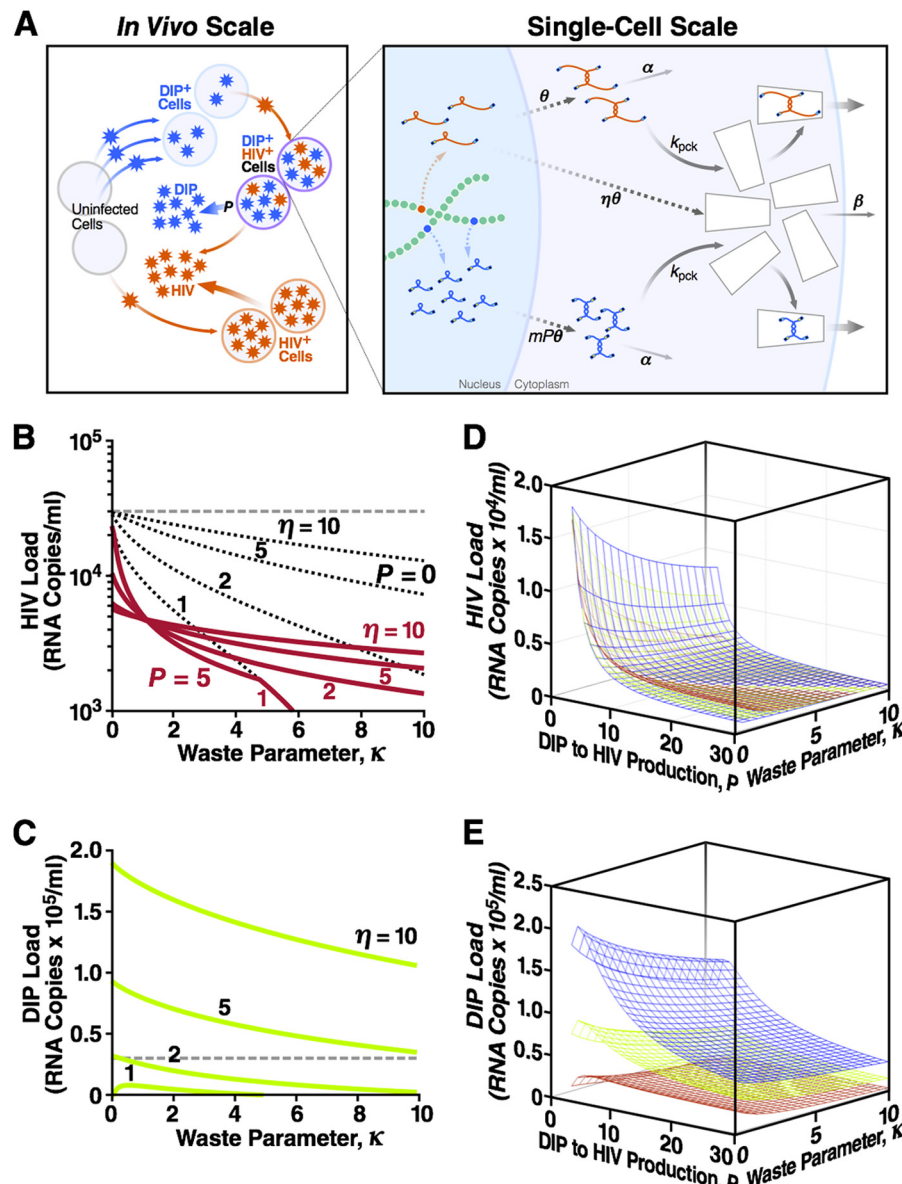
**FIG 1** Divergent evolution of the HIV-1 and DIP dimerization initiation sequences (DIS) by double mutations in HIV-1 indicates that DIP interference by genome stealing is evolutionarily unstable. (A) Genomic RNA (gRNA) monomers of HIV-1 and a DIP form three types of dimer complexes (HIV-HIV, HIV-DIP, and DIP-DIP) based upon a “kissing-loop” formation between the dimerization initiation sequences of HIV-1 and DIP, which contain a palindromic sequence (e.g., the consensus sequence GCGCGC). Due to a higher rate of transcription and multiple provirus copies, DIP monomers are more abundant, so most HIV-1 RNA is wasted on nonviable HIV-DIP heterodimers. (B) A simplified model representing the abundance of gRNA monomers for HIV-1 and a DIP in the cytoplasm of the infected cell:  $g(t)$  and  $g_{\text{DIP}}(t)$ , respectively.  $\theta$  is a lumped parameter representing the linear rate of gRNA production, and  $P$  is the expression asymmetry between HIV-1 and DIP.  $k_H$ ,  $k_{\text{DIP}}$ , and  $k_{\text{HIP}}$  are dimerization coefficients for HIV-HIV, DIP-DIP, and HIV-DIP, respectively. (C) Potential mutations in the kissing loop lead to divergent evolution of HIV-1 and DIP. Top row, in the wild-type (wt) HIV-1 and DIP case, there is an exact match for any gRNA pair (HIV-HIV, HIV-DIP, and DIP-DIP), which is enumerated in the rightmost column. Middle rows, if a single mutation arises within HIV-1 (highlighted by the blue rectangle), HIV-HIV homodimers have two mismatches, whereas HIV-DIP heterodimers have only a single mismatch and DIP-DIP homodimers have no mismatches. Bottom rows, in the (likely) scenario where the second compensatory mutation occurs in HIV, heterodimerization is disfavored compared to homodimerization. (D) Evolutionary fitness of homodimers and heterodimers qualitatively estimated based on dimerization coefficients in panel B and the number of sequence matches in panel C. In this idealized model, fitness takes the canonical functional form of an exponential where the selection coefficient,  $s$ , is based only upon the dimerization coefficients and the degree of sequence matching. Although a single mutation in the DIS of HIV-1 is more deleterious to HIV-1 homodimerization than to DIP-to-HIV-1 heterodimerization, a second mutation within the DIS will rescue HIV-1 dimerization and generate a further decrease in DIP-to-HIV-1 genome stealing.

and dimerized genomes (Fig. 1A and B) is used to describe gRNA pairing and to analyze how the gRNA pairing would coevolve for a DIP and HIV-1. The model describes the homozygous pairing of HIV-1 genomes ( $g$ ) and DIP genomes ( $g_{\text{DIP}}$ ) as well as the heterozygous pairing between DIP genomes and HIV-1 in dually infected cells. The model also captures experimental evidence demonstrating that subgenomic RNAs that share the consensus DIS palindrome of HIV-1 can dimerize to gRNA HIV-1 genomes (57, 58) and that partitioning of diploid genomes between homozygous and heterozygous virions is binomial (58). Importantly, the heterozygous virions that contain one copy of the HIV-1 genome and one copy of the DIP genome (subgenomic) are largely nonviable (59).

Furthermore, the model includes recent evidence that the overwhelming majority of HIV-1-infected cells harbor a single integrated HIV-1 provirus (60), most likely, due to the short lifetime of infected cells (61, 62) and molecular restrictions to superinfection, such as *nef* downregulation of surface CD4 (61). However, the model does not restrict DIP proviral integrations to a single copy, since multiple infections of the cell require expression of *trans* elements, which the DIP would lack. Here, specifically, we

consider a DIP that does not express Nef or any other *trans* elements responsible for protection against superinfection. Multiple copies of integrated DIP provirus, whose number we denote  $m$  ( $m = 1, 2, 3, \dots$ ), lead to the DIP gRNA monomers being more abundant than the HIV-1 gRNA in the cytoplasm of the cell by a factor of  $m$ , which varies among cells and is determined in part by the abundance of circulating virus in the body (i.e., viral dynamics). In addition, we assume that DIP genome architecture (e.g., the lack of splicing sites [63, 64]), leads to an expression asymmetry between a DIP and HIV, such that DIP monomers are more abundant in the cytoplasm of the cell by an additional fixed factor, which we denote  $P$  (where  $P > 1$ ). This “expression asymmetry” has been observed for lentiviral vectors (25, 59, 63–65), including HIV-1-derived lentiviral vectors (25, 59). The enrichment of the DIP gRNA over HIV gRNA in the cytoplasm is a product of both  $m$  and  $P$ . Due to enrichment of the DIP, most HIV-1 genomic RNA copies in the cell are stolen by the DIP to produce nonviable virions with HIV-DIP heterodimers, thereby generating a mechanism of interference.

To examine the evolutionary stability of interference by genome stealing (i.e., heterodimerization), we first consider



**FIG 2** DIPs that steal capsid proteins stably suppress the HIV-1 load across a broad range of parameters. (A) The model comprises two scales of biological organization. The *in vivo* (individual host) scale is the standard model of HIV-1 replication, expanded to include DIPs (see Supplemental Methods, equations S28 to S33, in the supplemental material). Uninfected cells can be infected with either HIV-1 or DIP, and DIP<sup>+</sup> cells can be superinfected with HIV-1 to become dually infected cells. The single-cell model is described by equations 1 to 3. A dually infected cell has one integrated HIV-1 provirus and multiple,  $m$ , copies of DIP provirus. A fraction of HIV-1 gRNA is translated into proteins that form “empty” capsids. The DIP does not express proteins. Dashed arrows represent multistage processes (including the loss of RNA monomers and capsid proteins). A fraction of stable dimer genomes and full capsids is also lost. Remaining genomes, HIV-1 or the DIP, are packaged within capsids and released as infectious particles. Shown also are the steady-state HIV-1 load (B) and the steady-state DIP load (C) at different values of two single-cell parameters: the capsid waste parameter,  $\kappa$ , and the capsid-to-genome production ratio,  $\eta$  (Table 1). The dashed line indicates the HIV-1 viral load in the absence of capsid waste and the DIP ( $\kappa = P = 0$ ), which is assumed to be the average load in untreated humans ( $3 \times 10^4$  RNA copies/ml blood). Calculations use a DIP/HIV-1 production ratio (i.e., expression asymmetry) of  $P = 5$  and a basic reproduction ratio of  $R_0 = 10$  (Table 1). The decrease in HIV-1 load in the presence of capsid waste ( $\kappa > 0$ , red lines), compared to the untreated HIV-1 “set-point” level (dashed line), is partly due to the loss of HIV-1 products (black dotted lines calculated at  $P = 0$ ) and partly due to the DIP, which competes with HIV-1 for available target cells and steals HIV-1 capsid in dually infected cells. The first effect is more important at  $\eta = \sim 1$ , and the DIP suppression factor is stronger at a large  $\eta$  value (see Fig. S2 in the supplemental material). Shown also are the steady-state HIV-1 load (D) and the steady-state DIP load (E) as functions of both expression asymmetry,  $P$ , and capsid waste parameter,  $\kappa$ , at three values of the capsid-to-genome ratio:  $\eta = 2$  (red),  $\eta = 5$  (green), and  $\eta = 10$  (blue). These 3D plots act as a partial sensitivity analysis showing that HIV-1 and DIP loads depend strongly on the value of  $P$ .

single-residue mutations in the HIV-1 DIS (e.g., GCGCGC → GCGAGC). Such mutations lead to mismatches and decreases in the probability of heterodimerization but result in more severe decreases in the levels of HIV-1 homodimerization (Fig. 1C).

However, compensatory double-residue mutations reestablish a new DIS palindrome (e.g., GCGCGC → GCUAGC) and generate a situation in which DIP-to-HIV-1 heterozygous dimerization is far less favorable than homozygous dimerization (Fig. 1C). By this

**TABLE 1** State variables and model parameters for the intracellular capsid-stealing model<sup>a</sup>

Notation	Definition	Unit	Value	Reference(s)
	State variables			
$G$	Concentration of full-length dimerized HIV genomic mRNAs	1/ $\mu$ l		
$G_{DIP}$	Concentration of full-length dimerized DIP genomic mRNAs	1/ $\mu$ l		
$C$	Concentration of encapsidation-competent capsids in cytoplasm	1/ $\mu$ l		
$m$	DIP (integrated) provirus copy number (i.e., MOI)	Dimensionless		
	Model parameters			
$\theta$	Rate of accumulation of encapsidation-competent HIV genomes in cytoplasm	1/ $\mu$ l/day		
$k_{pck}$	Packaging constant	$\mu$ l/day		
$\alpha$	Loss of genome rate	1/day		
$\beta$	Loss of capsid rate	1/day		
$\eta$	Capsid-to-genome accumulation ratio	Dimensionless	1.2 to 5	10 and 72
$P$	Expression asymmetry between DIP and HIV	Dimensionless	8 to 10	13

<sup>a</sup> See Fig. 2A and equations 1 to 3.

logic, the idealized model that considers dimerization coefficients to be a function of only the number of mismatches leads to divergent evolution of the DIS sequences between HIV-1 and a DIP due to double mutations in HIV-1 DIS (Fig. 1D). Although a single mutation in HIV-1's DIS is more deleterious to HIV-1 homodimerization than to DIP-to-HIV-1 heterodimerization, a second mutation within the DIS will rescue HIV-1 dimerization and generate a further decrease in DIP-to-HIV-1 genome stealing.

This analysis does not explicitly account for the cost of the palindromic DIS being different than the consensus GCGCGC sequence, since different HIV-1 strains encode different DIS palindromes (57). Another limitation of this analysis is that it only considers a single round of mutation. However, these assumptions are not critical for the conclusion of evolutionary instability of genome stealing. More realistic models that capture multiple rounds of mutation still lead to progressive decreases in heterodimerization levels relative to those of homodimerization (see Fig. S1 in the supplemental material).

**An alternative mechanism of interference: capsid stealing by DIPs.** Given that interference by genome stealing is unstable (see above), we next considered whether interference by DIP competition for *trans* elements, such as the capsid, was evolutionarily stable. Since DIPs carry the full complement of *cis* elements but lack one or more *trans* elements (e.g., capsid, envelope), DIPs must rely on the *trans* elements of the wild-type lentivirus to package and mobilize out of the infected cell. Thus, DIPs compete for and parasitize *trans* elements, leading to interference with the wild-type virus.

To calculate DIP interference on HIV-1 viral loads and the associated evolutionary stability, we used a recently developed multiscale modeling approach (17) to integrate a single-cell model of competition for capsid with an individual-patient model. Unlike previous multiscale modeling of virus infections (66–68), this multiscale model is designed specifically for HIV-1 in the presence of a DIP. The model applies to a range of DIP vectors, from a minimal DIP that does not code for any *trans* elements to a DIP that codes for *tat* and *rev* (allowing for the expression and export of genomes to the cytoplasm even in the absence of an HIV-1 provirus), but does not express capsid proteins or Nef (which mediates protection against superinfection). In the following subsections, we describe the single-cell and individual-host models, their integration, and their parameters. In the last subsection, we

introduce the notion of an effective selection coefficient. Detailed analytical derivations are presented in Supplemental Methods in the supplemental material. Numerical solutions for all figures were performed in MATLAB (version R2011a).

**The single-cell model with capsid stealing.** For tractability, the single-cell model considered here (Fig. 2A) is simplified and considers only intracellular replication events relating to dimerized wild-type HIV-1 RNA genomes ( $G$ ), encapsidation-competent capsid ( $C$ ), and dimerized DIP RNA genomes ( $G_{DIP}$ ). The equations have the following form:

$$\frac{dG}{dt} = \underbrace{\theta}_{\substack{\text{HIV genome} \\ \text{production}}} - \underbrace{k_{pck}GC}_{\substack{\text{Packaging of} \\ \text{HIV genomes}}} - \underbrace{\alpha G}_{\substack{\text{Loss of} \\ \text{HIV genomes} \\ \text{into capsids}}} \quad (1)$$

$$\frac{dC}{dt} = \underbrace{\eta\theta}_{\substack{\text{Capsid} \\ \text{production}}} - \underbrace{k_{pck}(G + G_{DIP})C}_{\substack{\text{Encapsulation of genomic RNAs}}} - \underbrace{\beta C}_{\substack{\text{Loss of} \\ \text{capsids}}} \quad (2)$$

$$\frac{dG_{DIP}}{dt} = \underbrace{mP\theta}_{\substack{\text{DIP genome} \\ \text{production}}} - \underbrace{k_{pck}G_{DIP}C}_{\substack{\text{Packaging of} \\ \text{DIP genomes}}} - \underbrace{\alpha G_{DIP}}_{\substack{\text{Loss of} \\ \text{DIP genomes} \\ \text{into capsids}}} \quad (3)$$

The model parameters are defined in Table 1. Briefly, the model describes the production and decay of dimerized HIV-1 genomes, the packaging of these dimerized genomes into capsids that are produced at a rate proportional to that of genome dimers, and the competition for encapsidation between DIP genomes and HIV-1 genomes. The model neglects heterozygous genomes since, as demonstrated above, the dimerization initiation sequence for HIV-1 and the DIP will diverge so that heterozygous pairing is eliminated. As in the genome-stealing model (above), we allow each DIP provirus to express more RNA than an HIV-1 provirus by a fixed factor of  $P$  (25, 59, 63–65). This expression asymmetry ( $P > 1$ ) is due to differences in genome architecture, such as mutated alternative splicing sites (63, 64). We classify dually infected cells by the number of DIP provirus copies  $m$  ( $m = 1, 2, 3, \dots$ ). The net asymmetry between a DIP and HIV-1 expression in a dually infected cell is the product  $m \times P$ . Based on previous results (17), we assume that the HIV-1 and DIP levels inside of the cells

TABLE 2 State variables and parameters for the individual-host model<sup>a,b</sup>

Notation	Definition	Unit	Value	Reference or source
State variables				
$T$	Uninfected CD4 <sup>+</sup> T cells permissive for viral replication	Cells/ $\mu$ l		
$I$	CD4 <sup>+</sup> T cells infected with HIV only	Cells/ $\mu$ l		
$T_{DIP,m}$	CD4 <sup>+</sup> T cells infected with $m$ copies of DIP provirus but not infected with HIV ( $T_{DIP,0} = T$ )	Cells/ $\mu$ l		
$I_{D,m}$	Dually infected cells with an HIV and $m$ copies of DIP provirus	Cells/ $\mu$ l		
$V$	HIV viral load	RNA copies/ml		
$V_D$	DIP viral load	RNA copies/ml		
Model parameters				
$b$	Linear production rate of uninfected cells	cells/ $\mu$ l/day		
$d$	Death rate of uninfected cells	1/day		
$k$	Infectivity factor	ml/day/RNA copy		
$n$	HIV burst size from a singly infected cell	RNA copy/cell		
$c$	Virion clearance rate	1/day		
$\delta$	Death rate of HIV infected cells	1/day	1.0/day	50
$m\psi_m$	HIV burst size from a dually infected cell with $m$ copies of DIP provirus	Dimensionless	$k_{pck}GC/\delta$	From single-cell model
$n\psi_m\psi_m$	DIP burst size from a dually infected cell with $m$ copies of DIP provirus	Dimensionless	$k_{pck}G_{DIP}C/\delta$	From single-cell model

<sup>a</sup> See equations 4 to 9.<sup>b</sup> In the two bottom rows,  $G$ ,  $G_{DIP}$ , and  $C$  are steady-state values of state variables for the intracellular model (see equations 1 to 3 and Table 1) and are derived in Supplemental Methods, equations S6 to S20, in the supplemental material.

rapidly reach a steady state; the steady-state values of  $G$ ,  $G_{DIP}$ , and  $C$  are derived analytically in Supplemental Methods in the supplemental material (see equations S6 to S14).

**The individual-host model.** Similar to the method previously used (17), the output of the single-cell model is used to calculate HIV-1 and DIP viral loads within an individual patient using a standard model of HIV-1 *in vivo* dynamics (55, 61, 69) that is generalized to include the production of DIPs. The generalized model includes coinfection of cells with DIPs and HIV-1 so that the dually infected cells produce less HIV-1. Based on the results of recent *in vivo* studies (60), we assume a single HIV-1 provirus per (singly or dually) infected cell but allow for multiple DIP provirus copies per cell, as previously proposed (35). Briefly, the model describes uninfected CD4<sup>+</sup> T cells permissive for viral replication ( $T$ ), cells infected with HIV-1 only ( $I$ ), CD4<sup>+</sup> T cells harboring  $m$  copies of DIP provirus but not infected with HIV-1 ( $T_{DIP,m}$ ) (by definition,  $T_{DIP,0} = T$ ), dually infected cells harboring a copy of HIV-1 and  $m$  copies of DIP provirus ( $I_{D,m}$ ), HIV-1 load ( $V$ ) (free virus concentration in peripheral blood plasma), and DIP load ( $V_D$ ). The system of equations has the following form:

$$\frac{dT}{dt} = b - (d + kV + kV_D)T \quad (4)$$

$$\frac{dI}{dt} = kVT - \delta I \quad (5)$$

$$\frac{dT_{DIP,m}}{dt} = kV_D T_{DIP,m-1} - (d + kV + kV_D)T_{DIP,m}, \quad m = 1, 2, 3, \dots \quad (6)$$

$$\frac{dI_{D,m}}{dt} = kVT_{DIP,m} - \delta I_{D,m}, \quad m = 1, 2, 3, \dots \quad (7)$$

$$\frac{dV}{dt} = n\delta I + n\delta \sum_{m=1}^{\infty} \psi_m I_{D,m} - cV \quad (8)$$

$$\frac{dV_D}{dt} = n\delta I + n\delta \sum_{m=1}^{\infty} \psi_m I_{D,m} - cV_D \quad (9)$$

The model parameters, which are well described in the literature and summarized in Table 2, are the linear production rate of uninfected cells ( $b$ ), the natural death rate of uninfected cells ( $d$ ), the infectivity factor ( $k$ ), the death rate of singly and dually infected cells ( $\delta$ ), and the HIV-1 burst size from a singly infected cell ( $n$ ). There are two additional parameters in the presence of a DIP:  $\psi_m$ , the ratio of HIV-1 burst size between a singly infected cell,  $I$ , and a dually infected cell with  $m$  copies of DIP provirus  $I_{D,m}$ , and  $\rho_m$ , the ratio of the DIP to HIV-1 burst size from a dually infected cell with  $m$  copies of DIP provirus.

The steady states of the single-cell model (equations 1 to 3) define the following burst sizes for HIV-1 and a DIP (see equations S6 to S20 in Supplementary Methods in the supplemental material):

$$n = k_{pck}[GC]_{p=0}/\delta \quad (10)$$

$$\psi_m n = k_{pck}GC/\delta \quad (11)$$

$$\rho_m \psi_m n = k_{pck}G_{DIP}C/\delta \quad (12)$$

These expressions serve as input parameters for the individual-host model. The individual-host model (equations 4 to 9) is similar to the model described by Metzger et al. (17), except that here we relax the restriction of a single DIP copy per cell and allow cells to have multiple DIP infections. We assume that the state variables of the individual-host model (equations 4 to 9) are in a steady state, which corresponds to chronic infection. Equations 4 to 9 are used to calculate steady-state levels, as described in equations S34 to S43 and the following subsection in Supplementary Methods in the supplemental material.

**Parameter values.** The full list of model parameters is given in Tables 1 and 2. Using a standard approach, we reduce the number of parameters to a smaller number of composite parameters by changing units to those in which the state variables are measured (see Supplemental Methods in the supplemental material). As a result of

this nondimensionalization, all results can be conveniently expressed in terms of two nondimensional parameters that capture the evolutionary potential of HIV-1: (i) the composite “waste” parameter  $\kappa = (\alpha\beta)/(\theta k_{pk})$ , which reflects the loss of HIV-1 genomes (rate  $\alpha$ , equation 1) and capsids (rate  $\beta$ , equation 2), and (ii) the ratio of the encapsidation-competent capsids to the dimerized HIV-1 genomes produced per unit time ( $\eta$ ) (referred to as the capsid-to-genome production ratio). The remaining parameters are determined from the basic reproductive ratio ( $R_0$ ), which is estimated from *in vivo* data as  $R_0 \sim 10$  (the case of an exponentially distributed production delay with an average between 24 h and 12 h) (see reference 70) (Table 3), and by using fixed values of  $P$ . Lentiviral DIPs with  $P$  values of 8 to 10 have been engineered (65). We focus on the conservative value of  $P = 5$  below, but the interval  $P = 2$  to 30 is also studied (see Supplemental Methods).

**Testing evolutionary stability: the effective selection coefficient.** To determine if HIV-1 mutates to increase or decrease capsid waste (or the capsid-to-genome rate ratio), we capitalized on previous studies that applied concepts from Darwinian evolution, based on the notion of the selection coefficient ( $s$ ), to analyze the evolution of single-locus mutations in the HIV-1 genome (41, 48). Negative values of  $s$  denote that a mutation decreases the progeny of an infected cell and is selected against, and positive values of  $s$  denote that a mutation is selected for. Calculating  $s$  is complicated by the compartmental structure of a population of infected cells that includes both dually ( $HIV^+ DIP^+$ ) and singly ( $HIV^+ DIP^-$ ) infected cells that may contribute differently to the fitness effect of mutation (number of progeny). The calculation of  $s$  is also affected by the dynamic interaction of HIV-1 and DIP within dually infected cells. To account for these complexities, we introduce the effective selection coefficient ( $\partial s_{eff}$ ), defined as the exponential rate of increase or decrease of a mutant virus normalized to the death rate of infected cells. To calculate  $\partial s_{eff}$  we begin with a DIP–HIV-1 coinfection at steady state and perturb the system by adding a small amount of mutant HIV-1. In particular, we consider mutations that increase capsid waste ( $\kappa \rightarrow \kappa + \partial\kappa, \partial\kappa > 0$ ), achieved by slightly reducing the value of  $k_{pkg}$  (Fig. 3A). We then calculate the expansion (contraction) rate of the mutant subpopulation from the model equations (equations 1 to 12) and arrive at normalized value  $\partial s_{eff}/(\partial\kappa/\kappa)$ . The normalized selection coefficient for mutations affecting the capsid-to-genome production ratio  $\eta$  is calculated in analogous fashion (detailed derivations are given in Supplemental Methods, equations S48 to S57, in the supplemental material).

## RESULTS

**Evolutionary stability of capsid-stealing interference: general approach.** Our preliminary analysis (see “Preliminary analysis”) showed that the genome-stealing mechanism of DIP interference with HIV-1 replication is evolutionarily unstable due to divergence of the HIV-1 and DIP dimerization initiation sequences (Fig. 1). Here, we determine whether interference by DIP competition for *trans* elements, such as the capsid protein, leads to stable and sustained interference. We use our recently developed multi-scale modeling approach to integrate a single-cell model with an individual-patient model (17). The single-cell model (equations 1 to 3) captures the reported ability of minimal lentiviral vectors to express more RNA in the cytoplasm than HIV-1 (25, 59, 63–65), and this expression asymmetry is represented by the parameter  $P$  (where  $P$  is  $>1$ ). The output from this single-cell model (i.e., DIP

and HIV-1 burst sizes) is used as the input for the individual-patient model (equations 4 to 9) (Table 2)—a generalized form of the standard model of HIV-1 *in vivo* dynamics (55, 61, 69) that includes the production of DIP particles—to ultimately calculate HIV-1 and DIP viral loads within a patient (Fig. 2A). Detailed analytical derivations and numeric calculations are presented in Supplemental Methods in the supplemental material.

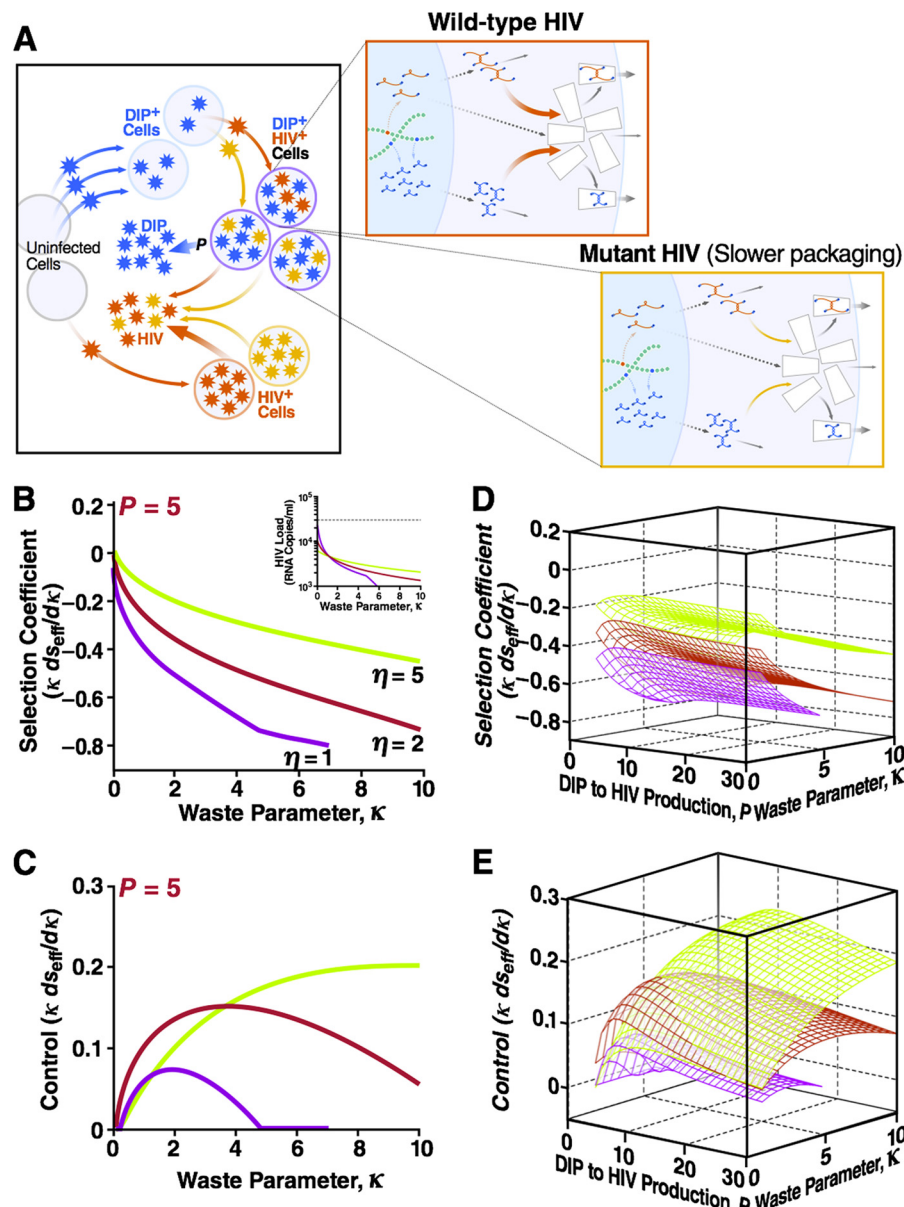
**DIP interference by competition for the capsid within single cells leads to sustained suppression of HIV-1 at the individual-patient level.** To test whether DIP lowers HIV-1 viral load, we examined the steady-state values of HIV-1 and the DIP at the individual-patient scale as a function of parameters in the single-cell model. As detailed above, the results are expressed in terms of a pair of intuitive and nondimensional parameters that capture the evolutionary potential of HIV-1 and the DIP: the composite waste parameter ( $\kappa$ ) and the ratio of encapsidation-competent capsids to dimerized HIV-1 genomes produced per unit time ( $\eta$ ).

Analytical solutions demonstrate that HIV-1 viral load is stably decreased across a broad range of  $\kappa$  values (Fig. 2B, red lines). In most cases, the viral load is decreased by more than 1 order of magnitude, which is associated with significantly reduced HIV-1 transmission and disease progression (71). Stable DIP-mediated suppression of HIV-1 depends upon the expression asymmetry ( $P > 1$ ) (17, 35) and, importantly, is amplified by the large average multiplicity of DIP infection (see next section; see also Fig. S3a in the supplemental material). However, the decrease in HIV-1 viral load is not completely due to the DIP and is partly due to the loss of HIV-1 capsids and genomes by increased capsid waste (black dotted lines). Nevertheless, DIP-mediated suppression contributes to a significant fraction of this decrease through the stealing of HIV-1 capsids within dually infected cells and through competition for available target cells within the infected individual (see Fig. S2 in the supplemental material). At high  $\kappa$  values, there is a loss of dynamic stability because at such high capsid waste, HIV-1 crosses the threshold of its own extinction.

Even at modest values of expression asymmetry ( $P$ ), the high multiplicity of DIP infection generates relatively high DIP viral loads (see next section; see also Fig. S3 in the supplemental material) and allows the DIP to be dynamically stable even when  $\eta$  is just greater than 1 (Fig. 2C).

Thus, DIPs that steal capsids can stably suppress HIV-1 viral load even if the expression asymmetry ( $P$ ) of the DIP is modest. These results are robust and qualitatively similar across a broad range of  $P$  and  $\eta$  values (Fig. 2D and E).

**Robust suppression of HIV-1 is due to a high multiplicity of DIP infection.** As previously noted (35), DIP-infected cells would not be restricted to harboring only a single DIP provirus, since DIP-infected cells are long lived and could be readily reinfected multiple times by the DIP before HIV-1 infects the cell. DIP superinfection will lead to multiple integrated DIP genomes per cell. The number of DIP copies (denoted  $m$ ) varies among dually infected cells as predicted by the individual-patient model (see equations 4 to 9). We calculated the average DIP copy number, denoted  $E[m]$ , as a function of capsid waste ( $\kappa$ ) and expression asymmetry ( $P$ ) for different values of the capsid-to-genome ratio ( $\eta$ ) (see Supplemental Methods and Fig. S3a in the supplemental material). The results are dependent upon  $P$  in a positive manner: increases in the value of  $P$  lead to more DIP virions, which results in a greater multiplicity of DIP infection. It is useful to note that, according to the standard individual-patient model we use here,



**FIG 3** A DIP-HIV interaction is evolutionarily stable over a broad parameter range: HIV-1 cannot escape DIP by decreasing packaging resources. (A) The two-scale model for an individual infected by two strains of HIV, wild type (red) and mutant (orange), as well as a DIP (blue). Mutation causes a small decrease in the packaging constant of both HIV-1 and DIP  $k_{pck}$  and, hence, an increase in capsid waste parameter,  $\kappa = \alpha\beta/(k_{pck})$ , when  $\partial\kappa > 0$ . (B) Normalized effective selection coefficient,  $\partial s_{eff}/(\partial\kappa/\kappa)$ , for that mutation as a function of  $\kappa$  for a range of capsid-to-genome production ratios  $\eta$ . Fixed parameters are as described for Fig. 2B and C:  $R_0 = 10$  and  $P = 5$ . The negative values of  $\partial s_{eff}/(\partial\kappa/\kappa)$  imply that the mutation has net deleterious effects on HIV-1 replication. Overall, HIV-1 mutations that increase capsid waste are selected against. Inset, HIV-1 load as a function of the waste parameter from Fig. 2B. (C) A negative control showing  $\partial s_{eff}/(\partial\kappa/\kappa)$  within HIV<sup>+</sup> DIP<sup>+</sup> dually infected cells when burst size changes due to increased capsid waste (the first term in equation S54 in the supplemental material) are neglected. Only in this specific context, when burst size changes are ignored, are HIV-1 mutations that increase capsid waste selected for. Shown also are the net  $\partial s_{eff}/(\partial\kappa/\kappa)$  (D) and control  $\partial s_{eff}/(\partial\kappa/\kappa)$  (i.e., when burst size changes are neglected) (E) as functions of both  $P$  and  $\kappa$ . These 3D plots act as a partial sensitivity analysis and show that the selection coefficient weakly depends on the value of  $P$ . Detailed calculations are given in Supplementary Methods in the supplemental material.

the probability that a DIP<sup>+</sup> HIV<sup>-</sup> cell is infected with another DIP copy is proportional to the concentration of DIP virions. Dually infected “producer” cells contain higher levels of DIP gRNA than HIV-1 gRNA (i.e.,  $P > 1$ ), and these cells generate more DIP than HIV-1 virions, which results in an increase of the average DIP copy number with  $P$ . Thus, the ratio of DIP/HIV-1 gRNA within a cell is determined by  $P$  in two ways: directly (through the molec-

ular architecture of the DIP) and indirectly through the increase in the multiplicity of DIP infection ( $m$ ).

To further explore the contribution of  $m$  to HIV-1 interference and suppression, we artificially limited DIP multiplicity to a single copy per cell (i.e.,  $m = 1$  or  $m = 0$ ) and recalculated the HIV-1 viral loads (see Fig. S3b in the supplemental material). This control demonstrates that when  $m \leq 1$ , DIP-mediated suppression contributes

very little to the decrease in the HIV-1 viral load (i.e., when  $m$  is  $\leq 1$ , HIV-1 suppression is modest and arises primarily due to increased capsid waste). The analysis further shows that when  $m$  is  $\leq 1$ , the DIP loses stability at low  $\eta$  values as  $\kappa$  increases. Hence, when  $m$  is  $\leq 1$ , a high  $\eta$  value is needed for even modest suppression of the HIV-1 viral load. In summary, the multiplicity of integrated DIP genomes ( $m$ ) is critical for DIP-mediated suppression of HIV-1.

**Capsid interference is evolutionarily stable.** We next examined the direction of the evolution of the DIP and HIV-1 to test whether the DIP would be selected against in an ongoing HIV-1 infection. Conceivably, HIV-1 could escape a DIP by mutating to effectively increase the capsid waste parameter ( $\kappa$ ). One possible mechanism for increasing  $\kappa$  is for HIV-1 to mutate its packaging signal  $\Psi$  and thus decrease the packaging efficiency, allowing more genomes, or capsids, to be degraded instead of packaged. Under this increased-waste scenario, capsid stealing by the DIP would be more affected than HIV-1 packaging in dually infected cells due to the DIP expression asymmetry ( $P > 1$ ) and integration multiplicity ( $m > 1$ ). Thus, increased capsid waste would benefit HIV-1 due to a decrease in DIP interference. However, in the absence of a DIP, mutation toward increased capsid waste would be deleterious for HIV-1, since an increased loss of capsid products lowers the HIV-1 burst size. With these competing pressures, it is not clear which evolutionary direction dominates. These competing effects of mutation in the packaging loop are in fact evident from the steady-state HIV-1 load versus  $\kappa$  (Fig. 2B), where there are two components of HIV suppression: one due to a decrease in HIV-1 burst (black dotted curves versus the dashed line in Fig. 2B) and another due to DIP interference (red curves versus black dotted curves in Fig. 2B). As  $\kappa$  is increased, one component becomes larger and another smaller. It is not immediately clear which effect is stronger.

To determine if HIV-1 mutates to increase or decrease capsid waste, we calculate the effective selection coefficient ( $\partial s_{\text{eff}}$ ), defined as the exponential rate of increase, or decrease, of mutant HIV-1 normalized to the death rate of infected cells (see Theory; see also Supplemental Methods, equations S48 to S57, in the supplemental material). Negative values of  $s_{\text{eff}}$  indicate that a mutation decreases the progeny of an infected cell and is selected against, while positive values of  $s_{\text{eff}}$  indicate that a mutation is selected for. We consider mutations that slightly increase the value of  $\kappa$  by reducing the packaging efficiency (Fig. 3A). Because the effect of a mutation on  $\kappa$ , denoted  $\partial \kappa$ , is unknown and may vary among bases, the selection coefficient is expressed in a normalized form:  $\partial s_{\text{eff}}/(\partial \kappa/\kappa)$ . Unlike previous work, this study is unique in that it calculates an *in vivo* selection coefficient value directly from a molecular model. Previous studies were only able to estimate  $s$  by fitting (40, 43, 46, 48, 52–54).

Analyzing the effective selection coefficient as a function of capsid waste ( $\kappa$ ) demonstrates that, overall, HIV-1 mutants with increased capsid waste are selected against, since  $\partial s_{\text{eff}}/(\partial \kappa/\kappa)$  is  $< 0$  for a range of  $\eta$  and  $\kappa$  values (Fig. 3B). Importantly, HIV-1 mutants with high capsid waste are selected against despite DIP interference being decreased at high capsid waste (see Fig. 2B for low values of  $\eta$ ).

As a negative control, we next examined the effective selection coefficient keeping the burst size of HIV-1 in singly infected cells constant as capsid waste increases. As expected, in this scenario HIV-1 does evolve toward high capsid waste, as shown by  $\partial s_{\text{eff}}/(\partial \kappa/\kappa)$

being  $> 0$  for a range of  $\eta$  and  $\kappa$  values (Fig. 3C). The direction of evolutionary selection appears robust across a broad range of  $P$  values (Fig. 3D and E). Hence, it is the decrease in HIV-1 burst size that causes the negative selection coefficient shown in Fig. 3B.

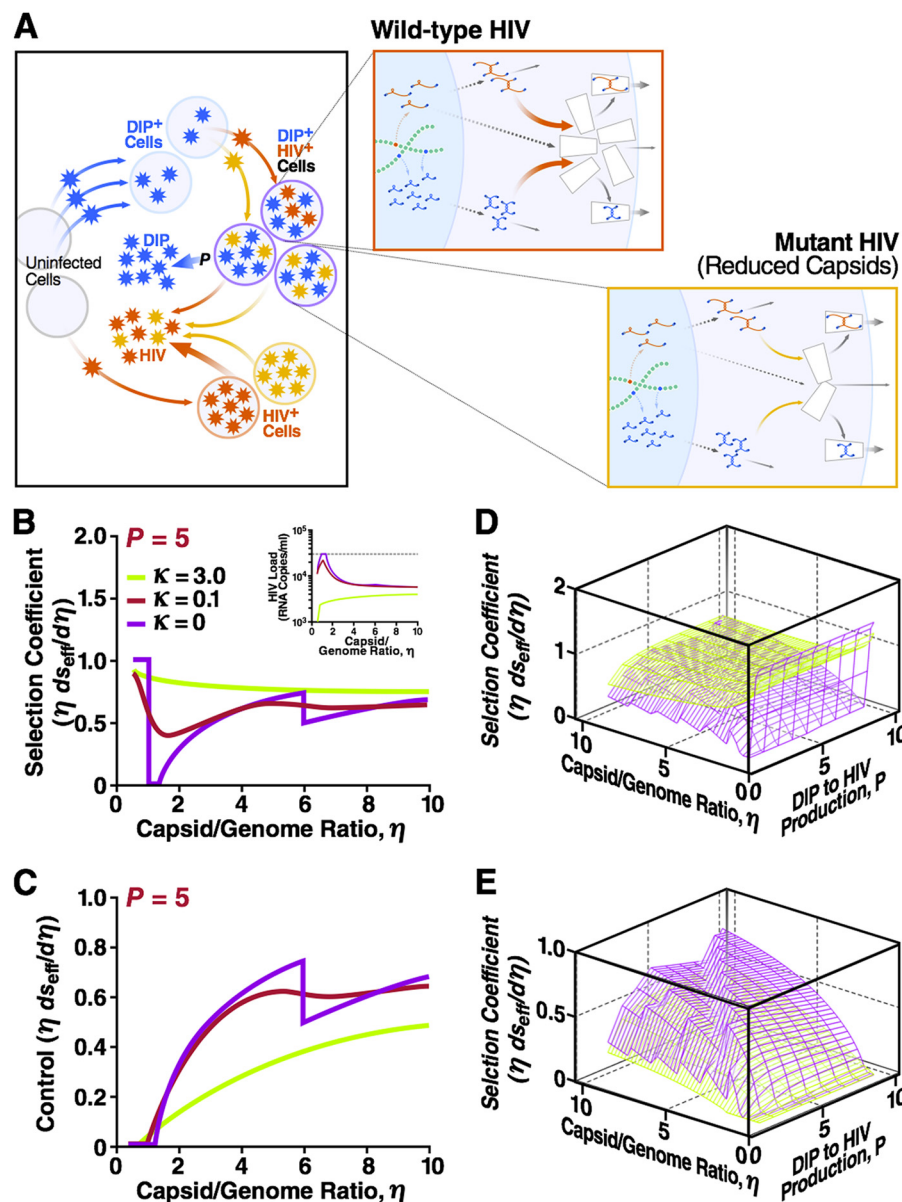
Essentially, these results indicate that the base HIV-1 burst size (which affects both singly and dually infected cells)—and not the DIP interference within dually infected cells—would dominate HIV-1 evolution within the individual patient.

Next, we sought to determine if HIV-1 could escape DIPs by mutating to reduce the available capsid material (i.e., decreasing its capsid-to-genome ratio  $\eta$ ) (Fig. 4A). The analysis shows that the DIP is unstable only at low values of  $\eta$ , and as expected, HIV-1 evolves toward high values of  $\eta$ , i.e., mutants that produce more capsid are selected for since  $\partial s_{\text{eff}}/(\partial \eta/\eta)$  is  $> 0$  for a large range of  $\kappa$  and  $\eta$  values (Fig. 4B). In fact, there is only a narrow band ( $1 < \eta < (P + 1)R_0/[P(R_0 - 1)]$ ) where the DIP is unstable and HIV-1 replication does not evolve toward increasing values of  $\eta$  (in this narrow band, the selection coefficient is zero). As a control, we examined keeping the HIV-1 burst size constant as the value of  $\eta$  increases (see the first term in equation S54 in the supplemental material). The control analysis shows that the effective selection coefficient is positive in the interval where the DIP is stable (Fig. 4C). Finally, a partial sensitivity analysis shows that the selection coefficient depends only weakly on the values of  $P$ , and the direction of evolutionary selection appears robust across a broad range of  $P$  values (Fig. 4D and E).

## DISCUSSION

This study examined the questions of whether lentiviral DIPs face evolutionary blocks and if DIPs would stably interfere with and suppress HIV-1. The analysis identified certain classes of DIPs that will not constitute an evolutionarily stable strategy of interference. Specifically, the analysis indicates that DIPs that suppress HIV replication via genome stealing will be unstable due to divergence between the HIV-1 DIS and the DIP DIS (i.e., interference and recombination via genome heterodimerization are minimized). This analytical finding raises doubts regarding the validity of a recent computational model that analyzes the evolutionary stability of conditionally replicating HIVs (i.e., DIPs) based exclusively on genome stealing (72). That DIS divergence is, in principle, possible is supported by the finding that different HIV-1 subtypes display diverse DIS sequences (73). Nevertheless, direct evidence demonstrating that the DIS is under selection from DIPs will require an intrapatient longitudinal data set showing evolution in the DIS.

As with all models, our analysis is a relatively simple representation of a complex system and necessarily makes certain assumptions. One potential caveat of our model is that it may not account for the finding that simple gammaretroviruses (not considered in the present work) exhibit spontaneous formation of DIPs (16), while lentiviruses do not. However, there are significant quantitative and qualitative differences in replication cycles between these two groups of viruses, and the model and parameters herein cannot be applied to the gammaretroviral life cycle. Most notably, HIV-1-infected cells are rapidly destroyed in a day or less, while in the gammaretrovirus setting, infected cells do not appear to display nearly as much cell death (except for mutants with poor protection against superinfection and an atypically broad host range) (74). Even more importantly, HIV-1 displays exclusively horizontal



**FIG 4** DIV-HIV interaction is evolutionarily stable over a broad parameter range. HIV-1 cannot escape a DIP by decreasing the capsid-to-genome ratio. (A) The two-scale model for an individual infected by two strains of HIV-1, wild-type (red) and mutant (orange), as well as a DIP (blue). Mutation causes a small increase in the capsid-to-genome ratio  $\eta$  by  $\partial\eta > 0$ . (B) Normalized effective selection coefficient  $\partial s_{eff}/(\partial\eta/\eta)$  for that mutation as a function of  $\eta$  for three values of the waste parameter  $\kappa$ . Fixed parameters used are as described for Fig. 2B and C:  $R_0 = 10$  and  $P = 5$ . Inset, corresponding HIV-1 viral load as a function of  $\eta$  at three values of the waste parameter  $\kappa$ . The positive values of  $\partial s_{eff}/(\partial\eta/\eta)$  imply that mutation is selected for, and HIV-1 evolves toward an increasing  $\eta$  value. When  $\kappa$  is 0 and  $\eta < 1$ , the DIP is not dynamically stable *in vivo*, and the selection coefficient is due exclusively to an increase in the HIV-1 burst size. In a narrow adjacent interval,  $1 < \eta < (P + 1)R_0/[P(R_0 - 1)]$ , DIP is still unstable, and HIV-1 replication does not require more capsid, which is why the selection coefficient is zero. (C) A negative control neglecting HIV-1 burst size changes due to mutation (the first term in equation S54 in the supplemental material) and showing that the effective selection coefficient is positive in the interval where the DIP is stable. The discontinuity at  $\kappa = 0$  and  $\eta = 1 + P$  is due to DIP gRNA competition with HIV gRNA for capsids at  $\eta < 1 + P$  but not at  $\eta > 1 + P$ , when there are enough capsids for both the DIP and HIV. Shown also are the net  $\partial s_{eff}/(\partial\eta/\eta)$  (D) and control  $\partial s_{eff}/(\partial\eta/\eta)$  (E) as functions of both  $P$  and  $\eta$ . Discontinuities at  $\eta = 1 + mP$ ,  $m = 1, 2, \dots$ , are analogous to the discontinuity described for panel C. These 3D plots act as a partial sensitivity analysis and show that the selection coefficient depends weakly on the value of  $P$ .

replication within a host (i.e., infection of new uninfected cells), which was taken into account in our model, while gammaretroviruses rely mostly on vertical replication (i.e., division of infected cells); their horizontal transmission is relatively infrequent and occurs between hosts or organs within a host. To determine whether these replication cycle differences account for the dis-

crepancy in DIP formation between gammaretroviruses and lentiviruses or whether additional factors are at play will require additional experimental and theoretical analysis. Additional simplifications of our model are designed to capture basic features of the lentiviral cycle inside of cells in conjunction with the within-host level. Specifically, we assumed that all process times are

Poisson distributed and that all dynamic variables are at steady state at both levels of modeling. A more complex model would include deterministic chaos that is present in all but the simplest dynamic systems, as well as various stochastic effects. For example, Kirkwood and Bangham (32) considered a generic virus and DIP coinfection in cell culture by introducing age-structured dynamics of infected cells and making the key assumption that wild-type virions are only produced from singly infected cells (i.e., perfect interference). In contrast, artificial lentiviral DIPs (25, 26, 29) do not exhibit perfect interference, and such DIPs would face strong negative selection pressures. Thus, it is unlikely that age-structured dynamics or time fluctuations would lead to chaotic dynamics for DIPs and HIV-1 or alter the main conclusions of our analysis, as long as the system remains far from instability.

Despite the caveats and assumptions inherent to any modeling exercise, the results herein provide a principled theoretical argument, based on the available data, that capsid stealing (competition for capsid proteins) is a dynamically and evolutionarily stable form of interference, provided that the ratio of capsid-to-dimerized genome production,  $\eta$ , is somewhat greater than 1. Capsid stealing is dynamically stable, primarily due to a multiplicity of DIP infection (i.e., multiple copies of integrated DIP provirus) that is higher than that of HIV-1, which appears to have a multiplicity of infection (MOI) of  $\approx 1$  (60). High MOIs are likely for DIPs that have deletions in transactivating elements, since these DIP-infected cells would not be subject to rapid virus-mediated or immune-mediated death; HIV-1-infected cells have an average life span of  $\sim 1$  day (61, 75) or less (76) and would therefore provide a long-lived target population for DIP superinfection. Also, cells infected with DIPs that lack *trans*-acting factors (such as HIV-1 Nef and Vpu) would not downregulate the cellular CD4 entry receptor that typically prevents HIV-1 superinfection of the cell (77).

The analysis also finds that capsid stealing is evolutionarily stable because the obvious mechanism that HIV-1 could use to escape DIP—increasing capsid waste ( $\kappa$ ) by decreasing packaging efficiency ( $k_{pck}$ )—ultimately harms HIV-1 replication significantly more than DIP production. HIV-1 could conceivably negate the deleterious effect of high capsid waste by generating compensatory mutations to ensure that DIP packaging is affected more than HIV-1 packaging (i.e., different  $k_{pck}$  in equations 1 and 3). However, to escape in this manner, HIV-1 would need to mutate both *cis* and *trans* elements (e.g., generate correlated double mutations in both the capsid protein and the packaging signal), while the DIP could counteract with a single mutation in its *cis* sequence. Thus, this escape approach would not benefit HIV-1 in the long run.

In principle, there are a number of possible scenarios for the value of  $\eta$  and its rate of evolution within an individual patient. The model predicts that effective selection coefficients for mutation in  $\eta$  are positive (Fig. 4). If  $\eta$  is  $<1$ , then the analysis above argues that HIV-1 will evolve to  $\eta$  equaling 1 rapidly (Fig. 4B), probably on the time scale of years, given the high rate of HIV-1 evolution. The evolution to  $\eta$  equaling 1 is predictable, since each genome requires a capsid. When  $\eta$  is  $>1$  in the absence of a DIP, HIV is still predicted to increase the value of  $\eta$  but at a much lower rate. The reason why HIV-1 evolves to a larger  $\eta$ , even when  $\eta$  is  $>1$ , is to compensate for capsid waste, but the rate of evolution in this regime is low because capsid waste is small (small  $\kappa$ ). Determining the rapidity of this evolution will be critical for determin-

ing when DIPs may arise and transmit in the population. Once  $\eta$  reaches the DIP stability threshold (slightly above  $\eta = 1$ ), HIV-1 evolution toward larger  $\eta$  values accelerates again due to the decrease of DIP interference (until  $\eta$  reaches a molecular limit) (Fig. 4B). The central findings of this analysis, the evolutionary instability of genome stealing and the evolutionary stability of capsid stealing when  $\eta$  is  $>1$ , conform to the conventional wisdom that a parasite (DIP) that harms its host (HIV-1) without need will be not be evolutionarily stable. Conversely, in the capsid-stealing scenario, a DIP that depletes a surplus of HIV-1 capsids is expected to survive.

While the value of  $\eta$  was not directly measured, analysis of the fraction of empty capsids versus capsids “filled” with RNA genomes can provide estimates of  $\eta$  (see Fig. S4 in the supplemental material). *In vivo* measurements (78, 79) suggest that the fraction of filled capsids is  $\sim 20\%$ , which implies  $\eta$  being  $\sim 5$  (80). While *in vitro* measurements in a tissue culture viral expression system (293T human embryonic kidney cells) estimated the fraction of filled capsids to be 90% (58), which sets a bound of  $\eta$  being  $\leq 1.1$ , it is not unlikely that viral production in this 293T tissue culture setting is optimized compared to the *in vivo* setting. Thus, direct empirical measurements of  $\eta$  should be carried across a large sample of patients, and theoretical studies of HIV-1 evolution in a population of animal hosts could be used to predict the values of  $\eta$  that could evolve in the natural host.

The DIP stability condition ( $\eta$  slightly above 1) hints at a hypothetical schematic for the apparent absence of lentiviral DIPs. For example, historically, HIV-1 (or SIV) variants limited to  $\eta$  being  $<1$  due to intrinsic molecular and cellular factors might have penetrated a population already harboring HIV-1 (or SIV) strains with  $\eta$  at  $>1$  and already coinfect with a DIP. This new penetrating strain (with  $\eta < 1$ ) would transmit more efficiently, since it would likely not be suppressed by DIPs, and could spread across the population, causing depletion of DIPs. However, as HIV-1 evolves toward large  $\eta$  values within patients and  $\eta$  exceeds the DIP stability threshold, DIPs could reemerge and spread in a population, causing another round of preferential spread of an HIV-1 strain with  $\eta$  being  $<1$ . This cycle could repeat and lead to repeated boom-then-bust cycles for the prevalence of DIPs and HIV-1 strains with  $\eta$  values of  $>1$  and  $<1$  over time. We emphasize that we know of no evidence to support this historical speculation, and systematic population level theoretical studies will be required to determine the likelihood of this scenario over others and to determine when DIPs may arise and transmit in the population, as has occurred for dengue virus (7).

Regarding the safety of DIPs as an antiviral strategy, there is a long-standing concern of the possibility of rescuing a virulent phenotype by recombination between DIPs and the full-length HIV-1. However, the analysis here argues that, for HIV-1, this recombination is unlikely, since recombination requires heterodimerization via the DIS, and the HIV-1 and DIP DIS sequences diverge. Moreover, production of such a recombinant, even if possible, would be equivalent to the production of an HIV strain with lower fitness than that of the wild type. Because this recombinant is selected against, in order to be significant, it would need to occur in each and every DIP-infected cell, which is highly unlikely.

To conclude, our analysis argues that DIPs would be evolutionarily stable within an individual, provided that the capsid-to-genome production ratio is sufficiently large. Our previous

analysis argued that a DIP would be dynamically stable in a population level infection of HIV-1 (17). However, to address whether DIPs would be evolutionarily stable at the population level, further multiscale analysis is needed, since complex coevolutionary dynamics will exist between HIV-1 and a DIP, and these dynamics will be compounded by the existence of multiple circulating HIV-1 strains encoding different levels of capsid production.

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